

**Report Submitted 9/10/2021 to the Vermont Office of the Attorney General for Introduction of a New Prescription Drug to Market  
Actelion Pharmaceuticals US, Inc.**

**Information required pursuant to 18 VSA § 4637(c),(d)**

**UPTRAVI® IV**

Requirement	Submission - UPTRAVI® IV
<p>18 VSA § 4637(c)(1) A description of the marketing and pricing plans used in the launch of the new drug in the United States and internationally</p>	<p>While specific marketing and pricing plans are not available in the public domain, generally we plan to market UPTRAVI® IV in the US and promote to appropriate healthcare professionals who treat PAH patients who are hospitalized and are temporarily unable to take their UPTRAVI® Tablets. The pricing plan has WAC set for 1800mcg vial at \$19,200.00 for a 30 day supply. The list price of UPTRAVI® IV is not reflective of discounts and rebates which may be available through Medicaid, Medicare, and commercial insurance. UPTRAVI® IV will also be discounted as required under the 340B program, Federal Supply Schedule, and other government programs. International approvals are pending.</p>
<p>18 VSA § 4637(c)(2) The estimated volume of patients who may be prescribed the drug</p>	<p>The estimated number of patients in the United States with a condition for which UPTRAVI® IV may be prescribed is not in the public domain or publicly available.</p>
<p>18 VSA § 4637(c)(3) Whether the drug was granted breakthrough therapy designation or priority review by the FDA prior to final approval</p>	<p>The drug was not granted breakthrough therapy designation and priority review.</p>
<p>18 VSA § 4637(c)(4) The date and price of acquisition if the drug was not developed by the manufacturer.</p>	<p>Not an acquisition</p>
<p>Note: as provided in 18 VSA § 4637(d), we are limiting the information reported to that which is in the public domain or publicly available</p>	